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The compressibility of cubic white, orthorhombic black and rhombohedral black phosphorus

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The effect of pressure on the crystal structure of white phosphorus has been studied up to 22.4 GPa. The α phase was found to transform into the α ' phase at 0.87 ± 0.04 GPa with a volume change of 0.1 ± 0.3 cc/mol. A fit of a second order Birch-Murghanan equation to the data gave $Vo = 16.94 \pm 0.08$ cc/mol and $K_o = 6.7 \pm 0.5$ GPa for the α phase and $Vo = 16.4 \pm 0.1$ cc/mol and $K_o = 9.1 \pm 0.3$ GPa for the α ' phase. The α ' phase was found to transform to the A17 phase of black phosphorus at 2.68 ± 0.34 GPa and then with increasing pressure to the A7 and then simple cubic phase of black phosphorus. A fit of a second order Birch-Murnaghan equation to our orthorhombic and rhombohedral black phosphorus data gave $Vo = 11.43 \pm 0.02$

cc/mol and K_o = 34.7 ± 0.5 GPa for the A17 phase and Vo = 9.62 ± 0.01 cc/mol and K_o = 65.0 ± 0.6 GPa for the A7 phase.

I. INTRODUCTION

Elemental phosphorus exists as at least three allotropes at ambient conditions including white, black, and amorphous red [1]. Additional allotropes such as crystalline red, gray, violet, and brown, have been made under non-ambient conditions and scarlet, which forms from white when exposed to x-ray radiation. Of these white phosphorus (wP) is the most reactive; igniting in the presence of oxygen. It is widely used in munitions designed for signaling, screening and incendiary purposes. It is also used in the production of phosphoric acid, fertilizers, food additives and cleaning compounds and small amounts are used in pesticides and fireworks. White phosphorus exists as the α form at ambient conditions. It consists of P₄ tetrahedra arranged in the body centered cubic α-manganese structure with reported lattice parameters of 18.8Å [2] and 18.51Å [3]. When wP was subjected to high pressure a discontinuous change in volume was observed at 0.8GPa⁴ which was interpreted as due to a phase transition to a new phase of wP. The transition pressure was found to fall on decreasing temperature [4]. These experiments involved compressing wP to a set pressure, cooling the sample to the temperature of liquid air (-185.7°C) and then allowing the sample to warm. The transition was shown by Bridgman as an arrest in the regular rise in temperature with time [4]. This study gave a transition temperature of approximately -80°C at ambient pressure. An NMR study of wP found that cooling at ambient pressure to -165°C resulted in a transformation to the γ phase [5]. On heating, a γ phase wP

sample was found to transform to the β phase at -115°C and then to the α phase at -77°C. An x-ray study by Simon et al. (1987) [6] found the β to α transition to be at -76.4°C. A later study by Simon et al. (1997) [7] found the beta to α transition occurred at -76.5°C and the γ to beta transition occurred at -175°C using differential thermal analysis and, from x-ray diffraction results, the α to β transition occurred at about -80°C and the γ to β transition occurred at nearly -105°C. A Raman study [8] found the β to α transition to occur at -78°C and the γ to β transition to transpire at -183°C. In all these studies, temperature dependent phase transitions on warming were initiated from the γ phase. The β phase has been found to have a triclinic structure analogous to that of γ -plutonium. The structure of the γ phase was determined to be monoclinic but close to orthorhombic [9]. The P₄ tetrahedra in both the alpha and beta structures show librational disorder while the arrangement in the γ phase is virtually static [9]. Both of the low temperature transitions were found to be sluggish.

X-ray studies of wP are difficult due to the material partially transforming to scarlet phosphorus when exposed to x-rays [10]. Perhaps for this reason no measurement of the compressibility of either the alpha or beta forms of white phosphorus has been reported. Here we present an x-ray study of the compressibility of white phosphorus and subsequent black phosphorus phases at ambient temperature.

II. EXPERIMENTAL

White phosphorus, in stick form, was obtained and cut under water to achieve approximately 50 micron dimensions and then loaded with a pressure manometer into a diamond anvil cell (DAC) and pressurized in a hydrostatic medium consisting of 16:4:1 ratio of methanol:ethanol:water. Using an analytically calibrated confocal micro-Raman spectroscopy system it was verified that no impurities (e.g., phosphoric acid) were present above the 10 ppm concentration level. The 220 μm diameter by 70 μm tall sample chamber consisted of two counter opposed type Ia 400 μm diameter culet diamonds and a rhenium gasket used to laterally confine samples. Pressure manometers consisted either of ruby [11] or NaCl. Pressure was determined before and after each x-ray measurement and the average is reported. The ruby scale has an error in the pressure of ±5%. When determining pressure using ruby the 488nm excitation laser fluence and exposure time were minimized in order to minimize photo-conversion to grey phosphorus. Data were collected on beamline 12.2.2 at the Advanced Photon Source [12,13]. This beamline is equipped with a set of brightness preserving optics that delivers about 2 x 10¹⁰ photons per second into a 10μm focal spot. An x-ray energy of 25keV was used. A MAR345 imaging plate detector was used to collect diffracted x-rays. We limited the exposure of the sample to x-rays to 60 seconds in order to minimize conversion of the white phosphorus to scarlet phosphorus. A sample was reloaded whenever any sign of conversion to scarlet phosphorus was detected. Approximately ten exposures could be collected using 25keV radiation before any significant coloration of the sample or development of amorphous-like scattering background was observed. Data were collected up to a pressure of 22.4GPa. Two dimensional diffraction patterns were radially integrated using the fit2d program [14]. The sample to detector distance was determined using

the LaB₆ standard. Peak positions were determined either by whole pattern fitting using the GSAS package [15,16] or by fitting individual peaks with a pseudo-Voigt function. Molar volumes were calculated from the lattice parameters and used to determine values for zero pressure volume and bulk modulus using the Birch-Murnaghan equation of state [17]. Molar volumes were calculated for the α phase assuming 232 phosphorus atoms per unit cell [6].

III. THE STABILITY AND EQUATIONS OF STATE OF WHITE PHOSPHORUS

Hydrostatically compressed white phosphorus was found to remain in the body centered cubic phase up to 2.3GPa (FIG. 1) when it transformed to orthorhombic black phosphorus. Unit cell parameters and molar volumes are listed in table 1. No sign of the triclinic β phase was detected in this pressure range. Close examination of the data revealed a small discontinuity between 0.83 and 0.91GPa which we attribute to the transition observed by Bridgeman [4]. Since the x-ray diffraction patterns clearly show no indication of the beta phase of white phosphorus we will refer to it as the α' phase. Taking the mid point of these two pressure points gives a transition pressure of 0.87 ± 0.04 GPa compared to a value of about 0.8 GPa previously observed [4]. A plot of normalized stress as a function of strain (an Ff analysis) [18] was used to determine the bulk modulus and the pressure derivative of the bulk modulus for all data collected below 0.87 GPa. An analysis of the contributions to the calculated pressure from the various orders of strain in the Birch-Murnaghan equation revealed that only the second order term is significant in the pressure stability range of white phosphorus [18]. Consequently only a second order equation was fit to the data. A bulk modulus of 6.7(5) GPa was found for the α phase (FIG. 2). Since no ambient pressure volume exists for the α ' phase we have to first determine the ambient pressure

volume from a plot of normalized stress as a function of effective strain (a Gg plot) [18] (FIG. 2). A plot of normalized stress as a function of strain was then used to determine the bulk modulus (FIG. 2.). An ambient pressure volume of 16.4 ± 0.1 cc/mol and a bulk modulus of 9.1 ± 0.3 GPa was found. Thus the transition at 0.87 GPa is accompanied by a 36% decrease in compressibility. A gG analysis was also carried out for the α phase data, which gave a value for the ambient pressure volume of 16.94 ± 0.08 cc/mol. This compares well with the literature values of 16.99 cc/mol [3] and 17.02 cc/mol [19] and hence was used in our Ff analysis to compute the bulk modulus. The ambient pressure volume that we determined gives an ambient pressure lattice parameter for the α phase of 18.68 ± 0.09 Å.

IV. THE STABILITY AND EQUATIONS OF STATE OF BLACK PHOSPHORUS

The α ' phase of white phosphorus transformed into the orthorhombic A17 phase of black phosphorus (bP) at 2.68 \pm 0.34 GPa. We further compressed the sample to a maximum pressure of 22.4 GPa. The sample converted from the orthorhombic to the rhombohedral (A7) and then the simple cubic (sc) phase of as expected from previous studies of pure black phosphorus [20,21]. On depressurization the sample was found to transform from the sc to the A7 and then at ca. 0.5 GPa, to the A17 phase and to remain in the A17 phase at ambient pressure. Values for lattice parameters and molar volumes are contained in table 1.

Figure 4 contains a comparison of our A17 phase molar volumes with those of Kikegawa and Iwasaki [20]. Our data are seen to be better constrained than the previous data although they do

not cover the pressure range as well. We combined our data together with that of Kikegawa and Iwasaki [20] and used Gg and Ff plots to determine the ambient pressure volume, bulk modulus and the pressure derivative of the bulk modulus. Only the second order term was found to be significant in the pressure stability range of orthorhombic black phosphorus but the results of fits to a second and third order Birch-Murnaghan equation are included to allow comparison with the study of Akahama et al. [21]. We determined a zero pressure volume (V_o) of 11.43 K_o 0.05 cc/mol and a bulk modulus (K_o) of 34.7 \pm 0.5 GPa for a fit of a second order Birch-Murnaghan equation and a V_o of 11.43 \pm 0.05 cc/mol, a K_o of 26.5 \pm 0.1 GPa and the pressure derivative of the bulk modulus (K_o) of 10 ± 1 for a third order Birch-Murnaghan equation.

A comparison of our data for rhombohedral black phosphorus and that of Kikegawa and Iwasaki¹⁸ is illustrated in figure 5. Again our data are better constrained and have an adequate coverage of the pressure range so we used our data to determine a zero pressure volume, bulk modulus and pressure derivative of the bulk modulus as above. Here we determined $V_o = 9.62 \pm 0.01$ cc/mol and $K_o = 65.0 \pm 0.6$ GPa with a second order Birch-Murghanan equation and $V_o = 9.62 \pm 0.03$ cc/mol and $K_o = 55 \pm 1$ GPa and $K_o' = 7.5 \pm 0.5$ for a third order equation.

We only collected three data points in the simple cubic phase. These are compared with the data of Kikegawa and Iwasaki [20] and Akahama et al. [21] in Figure 6. We combined our data with that of Kikegawa and Iwasaki [20] and determined $V_o = 9.23 \pm 0.01$ cc/mol and $K_o = 72.5 \pm 0.3$ GPa with a second order Birch-Murghanan equation and $V_o = 9.23 \pm 0.03$ cc/mol and $K_o = 78.6 \pm 0.7$ GPa and $K_o' = 3.24 \pm 0.05$ for a third order equation.

V. CONCLUSIONS

White phosphorus was found to transform from the initial body centered cubic α form to a less compressible body centered cubic form, called the α ' form, at 0.87 ± 0.04 GPa with a 3.5% reduction of molar volume. This transformation occurred at approximately the same pressure as that observed by Bridgeman but does not correspond to the α to β transition observed at -76.9 °C and thought to extend with increasing pressure and temperature to 1.2 GPa and 64.4 °C [3]. The α ' phase is seen to have a smaller zero pressure volume and larger bulk modulus, or incompressibility, than the α phase which we would expect. White phosphorus transformed into orthorhombic black phosphorus at 2.68 ± 0.34 GPa and then transformed into rhombohedral and simple cubic black phosphorus with increasing pressure as expected from previous work [20,21]. On decreasing pressure the sample remained as black phosphorus transforming from simple cubic to rhombohedral and then orthorhombic in which form it remained to ambient pressure. Our orthorhombic and rhombohedra data were used to determine ambient pressure volumes and bulk modulli. These results provide additional constraints to previously published values.

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Phase	Pressure (GPa)	a(Å)	b(Å)	c(Å)	Molar volume (cc/mol)
α-wp	0.06(10)	18.642(9)			16.816(8)
α-wp	0.14(10)	18.559(9)			16.593(8)
α-wp	0.13(10)	18.548(9)			16.563(8)
α-wp	0.31(10)	18.421(9)			16.225(8)
α-wp	0.52(10)	18.255(9)			15.791(8)
α-wp	0.72(10)	18.147(9)			15.512(8)
α-wp	0.83(10)	18.088(9)			15.361(8)
α'-wp	0.91(10)	18.021(9)			15.191(8)
α'-wp	0.98(10)	17.966(9)			15.052(7)
α'-wp	1.17(10)	17.862(9)			14.793(7)
α'-wp	1.19(10)	17.860(9)			14.788(7)
α'-wp	1.21(10)	17.832(9)			14.718(7)
α'-wp	1.66(8)	17.670(9)			14.321(7)
α'-wp	1.81(9)	17.630(9)			14.224(7)
α'-wp	2.23(11)	17.424(9)			13.731(7)
o-bp	2.58(12)	3.302(7)	9.94(2)	4.22(1)	10.43(7)
o-bp	3.8(1)	3.299(4)	9.935(9)	4.206(5)	10.38(3)
o-bp	4.1(1)	3.302(3)	9.919(6)	4.206(4)	10.37(3)
o-bp	4.5(1)	3.301(5)	9.922(9)	4.192(4)	10.34(3)
h-bp	7.3(3)	3.388(7)		8.669(2)	8.65(3)
h-bp	10.0(1)	3.374(9)		8.604(3)	8.51(4)
sc-bp	13.6(3)	2.375(2)			8.067(7)
sc-bp	17(1)	2.346(4)			7.78(1)
sc-bp	22.4(8)	2.321(2)			7.531(5)
h-bp	9.44(10)	3.393(9)		8.669(2)	8.67(4)
h-bp	7.5(1)	3.389(2)		8.709(1)	8.69(1)
h-bp	5.37(10)	3.414(3)		8.810(2)	8.93(2)
h-bp	4.6(1)	3.420(3)		8.857(1)	9.00(1)
h-bp	4.23(10)	3.423(3)		8.887(1)	9.05(1)
h-bp	3.1(1)	3.434(5)		9.010(2)	9.24(3)
h-bp	3.0(1)	3.434(3)		9.010(2)	9.24(2)
o-bp	2.1(3)	3.29(1)	10.11(2)	4.20(2)	10.5(1)
o-bp	0.0(3)	3.310(2)	10.479(1)	4.387(5)	11.45(2)

Table I. Lattice parameters and molar volumes determined for white phosphorus (wp) and subsequent black phosphorous (bp) phases. o-bp the orthorhombic phase of black phosphorus, h-bp the hexagonal phase of black phosphorus and sc-bp the simple cubic phase of black phosphorus.

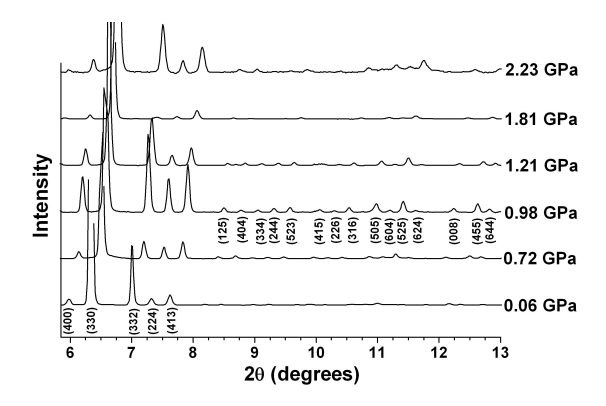


FIG. 1. Plot of a number of diffraction patterns of white phosphorus taken from ambient to just below the phase transition to orthorhombic black phosphorus. All of the data were not plotted for clarity. The miller indices for the strongest peaks are included.

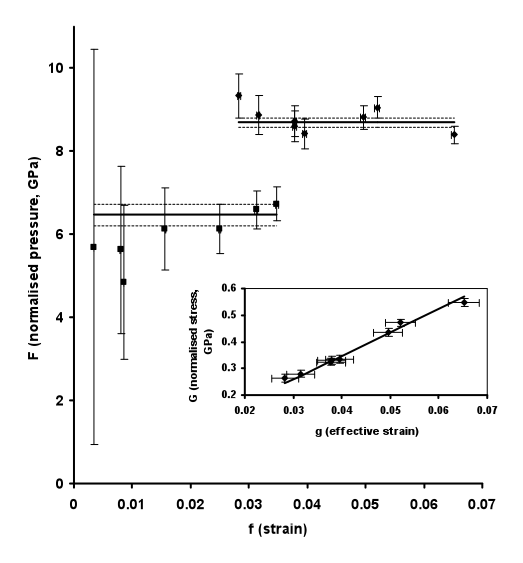


FIG. 2. The main figure shows plots of strain $(f=0.5x((V/V_0)^{-2/3}-1))$ against normalized pressure $(F=P/(1.5x((V/V_0)^{-7/3}-(V/V_0)^{-5/3})))$ for the α and α' phases of white phosphorus where P is the pressure (GPa), V is the molar volume at pressure P and V_0 is the molar volume at ambient pressure. The insert shows a gG plot $(g=0.5x((V^*/V)^{-2/3}-1), G=P/(3x(1+2g)^{3/2}))$ where P is the pressure (GPa), V is the molar volume and V^* an arbitrary reference molar volume) for the α' phase of white phosphorus.

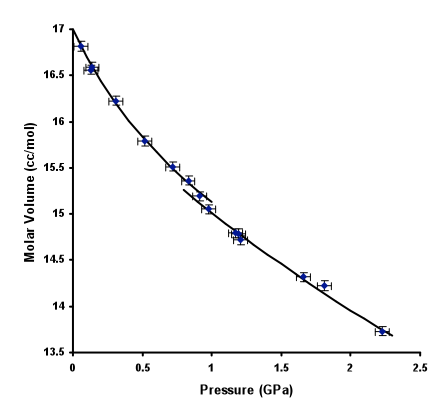


FIG. 3. The volume of white phosphorus as a function of pressure. The diamonds are the measured data points and the lines are fits of a second order Birch-Murghanan equation to the data above and below 0.87 GPa.

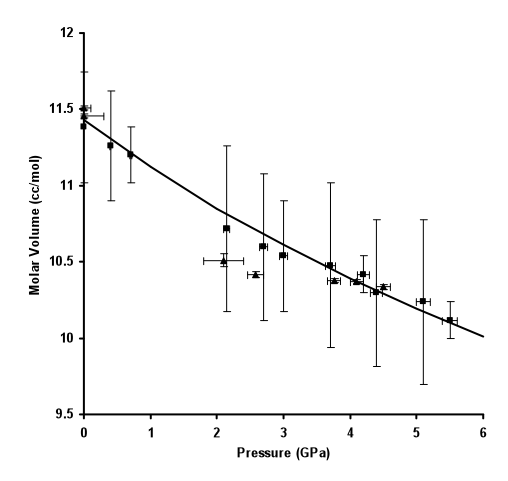


FIG. 4. The volume of the orthorhombic form of black phosphorus as a function of pressure. Our data are shown as filled triangles and the data of Kikegawa and Iwasaki [20] are shown as filled squares. A fit of a second order Birch-Murnaghan equation to all of the data is shown as a solid line.

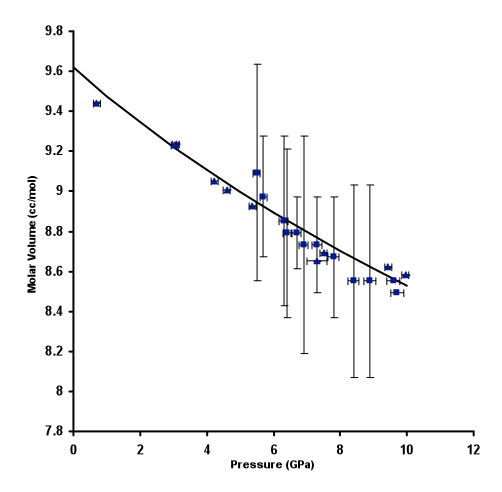


FIG. 5. The volume of the rhombohedral form of black phosphorus as a function of pressure. Our data are shown as filled triangles and the data of Kikegawa and Iwasaki [20] are shown as filled squares. A fit of a second order Birch-Murnaghan equation to all of the data is shown as a solid line.

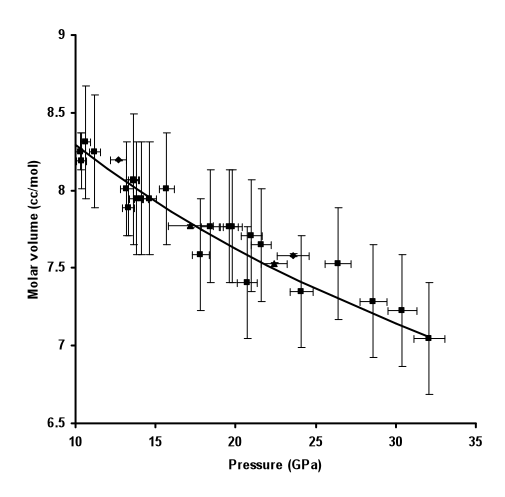


FIG. 6. The volume of the simple cubic form of black phosphorus as a function of pressure. Our data are shown as filled triangles the data of Kikegawa and Iwasaki [20] are shown as filled squares and the data of Akahama et al. [21] are shown as filled stars. A fit of a third order Birch-Murnaghan equation to our data and the Kikegawa and Iwasaki data is shown as a solid line.

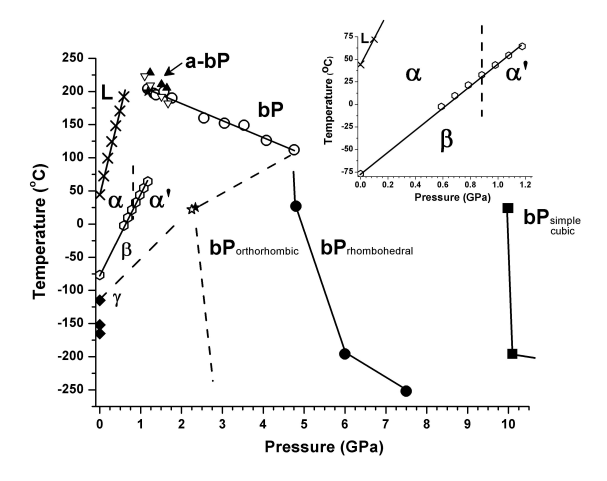


FIG. 7. A composite phase diagram of phosphorus including various forms that can originate from the white allotrope. The white allotrope stability field consists of the γ phase, solid diamonds at ambient pressure by Simon et al. [7] and Okudera et al. [9]; the β - α phase boundary, open hexagons by Bridgman [4]; the α - α phase boundary at ambient temperature, dashed vertical line at 0.87 GPa (see inset), this work; the solid alpha-liquid transition line, crosses by Bridgman [4]; the α -black phosphorus (bP) transitions, open circles by Patz [22], open downward triangles to amorphous black phosphorus (a-bP) (70

min. conversion) and solid upward triangles to a-bP (10 min. conversion) by Jacobs [23], black stars to a-bP and bP orthorhombic phases by Bridgman [4], α '-bP orthorhombic, open white star this work; the γ white-bP orthorhombic dashed transition line is drawn only as a suggestion; the bP orthorhombic-bP rhombohedral transition line, solid circles by Shirotani et al, [24]; and the bP rhombohedral-bP simple cubic transition line by Kikegawa [20].